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(54) Title: PULVERULENT MATERIALS

(57) Abstract: Pulverulent materials which contain surface-modified and structure-modified pyrogenically prepared metalloid or metallic oxide for the purposes of improvement.

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### Pulverulent materials

The present invention relates to pulverulent materials, to mixtures thereof, and to the use of surface- and structure-modified pyrogenically prepared metalloid or metallic oxides as flow regulators.

Pulverulent materials or their mixtures have a tendency to caking during prolonged storage.

It is known to add flow improvers or anticaking agents to pulverulent materials.

- 10 Known flow improvers are pyrogenic or precipitated, hydrophobic or hydrophilic silicas, silicates or metallic oxides. Pyrogenic silicas have higher purity compared with products prepared by precipitation. Particularly effective flow regulators (= synonymous with flow improvers or
- anticaking agents) are hydrophobic pyrogenic silicas, which are prepared by after-treatment of pyrogenic silicas with, for example, alkylsilanes or silicone oils. Known examples are AEROSIL R972, AEROSIL R974, AEROSIL R812 and AEROSIL R202 (manufactured by Degussa AG).
- Those known flow improvers and anticaking agents have the disadvantage that they do not always adequately improve the flowability. In addition, they are difficult to incorporate because they have a tendency to form agglomerates.
- If, for example, it is desired to use pyrogenic silicas as flow regulators, it is necessary to ensure that the agglomerates are comminuted mechanically to such an extent that the silica can develop its full effectiveness. This can be achieved by the following measures, for example:
- sieving the silica or the metallic oxide before it is
   mixed with one or more pulverulent materials,

- preparing a pre-mix of the pulverulent material and the silica or metallic oxide, which is then sieved,
- long mixing times and/or
- the use of mixing devices which exert sufficiently high shear energy to adequately comminute the agglomerates of the silica or of the metallic oxide.

That known process for the preparation of mixtures of pulverulent materials containing pyrogenic silicas or metallic oxides as flow regulators is accordingly relatively intensive in terms of time and energy. In addition, suitable devices, such as, for example, sieving machines or mixers, must be present.

The object of the present invention is to prepare

15 pulverulent materials and mixtures thereof which do not
have the disadvantages mentioned above, and to provide
pyrogenic silicas and metallic oxides as flow regulators
and anticaking agents in solids.

The invention provides pulverulent materials, and mixtures thereof, which are characterised in that they contain one or more pyrogenically prepared surface-modified and structure-modified metalloid or metallic oxides.

The invention also provides a method of improving the flowability of pulverulent materials and mixtures thereof, which method is characterised in that there are added to the pulverulent materials, or mixtures thereof, one or more pyrogenically prepared surface-modified, structure-modified metalloid or metallic oxides.

The invention relates also to the use of pyrogenically
prepared surface-modified and structure-modified metalloid
or metallic oxide for improving the flowability of

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pulverulent materials, pulverulent solids and mixtures thereof.

The pulverulent materials may consist of a single material or they may contain a mixture of various materials.

5 It has been found, surprisingly, that surface- and structure-modified pyrogenically prepared metalloid or metallic oxides are effective flow regulators and anticaking agents. They are easier to process and produce better flow behaviour than conventional pyrogenic silicas and metallic oxides.

According to the invention, the following pyrogenically prepared surface-modified and structure-modified metalloid or metallic oxides can be used:

(1) Silanised structure-modified silicas, which are characterised by groups attached to the surface, the groups being alkylsilyl (SiC<sub>n</sub>H<sub>2n+1</sub>, where n = from 2 to 18), preferably octylsilyl and/or hexadecylsilyl, having the following physicochemical data:

BET surface area m<sup>2</sup>/g: 25-400

20 average primary particle size nm: 5-50
pH value: 3-10
carbon content %: 0.1-25

DBP number %: The DBP number is at least 10 % smaller than the DBP number of the corresponding silanised silica without structure modification. With very pronounced structure modification, the structure may be so degraded that the DBP number can no longer be determined.

As the pyrogenically prepared silica used as the starting silica, a silica prepared by high-temperature hydrolysis from  $SiCl_4 + H_2$  and  $O_2$  may be used.

A silica prepared by high-temperature hydrolysis and having the following physicochemical data may especially be used.

	TO CULL							
	AEROSIL 90	130	150 150	AEROS1L 200	300	380	AEROSIL OX 50	TT 600
Behaviour towards water				hydro	hydrophilic			
Appearance				loose wh.	loose white powder			
BÉT surface area 1) m <sup>2</sup> /g	90 ± 15	130 ± 25	$150 \pm 15$	200 ± 25	300 # 30	380 ± 30	50 ± 15	200 ± 50
Average primary	20	16	14	12	7	6	40	40
particle size nm								
Tamped density $^2$ )				Ì				
normal product	ca. 80	ca. 50	ca. 50	ca. 50	ca. 50	ca. 50	ca. 130	ca. 60
T/6	l	Ca. 120	Ca. 120	Ca. 150	Ca. 120	Ca. 140	1	1
compressed product								
(addition of "V")			•					
Loss on drying 3)							]	
(2 hours at 1000°C) %	< 1.0	< 1.5	< 0.5 9)	<.1.5	< 1.5	< 1.5	< 1.5	< 2.5
on leaving the supplier								
Ignition loss $4)^7$ ) &	< 1	< 1	< 1	< 1	< 2	< 2.5	< 1	< 2.5
(2 hours at 1000°C)								
pH value 5) (in 4 % aqueous	3.6-4.5	3.6-4.3	3.6-4.3	3.6-4.3	3.6-4.3	3.6-4.3	3.8-4.8	3.6-4.5
dispersion)						•		•
	8 > 99.8	8.66 <	8.66 <	8.66 <	8.66 <	> 99.8	> 99.8	> 99.8
-	8 < 0.05	< 0.05	< 0.05	< 0.05	< 0.05	< 0.05	> 0.08	< 0.05
	8 < 0.003	< 0.003	< 0.003	< 0.003	< 0.003	< 0.003	< 0.01	< 0.003
	% < 0.03	< 0.03	< 0.03	< 0.03	< 0.03	< 0.03	< 0.03	< 0.03
	8 < 0.025	< 0.025	< 0.025	< 0:025	< 0.025	< 0.025	< 0.025	< 0.025
Sieve residue <sup>6</sup> ) &	< 0.05	< 0.05	< 0.05	< 0.05	< 0.05	< 0.05	< 0.2	< 0.05
(according to Mocker, 45 µm)								
(according to morker, 45 mil)								

- .1) following DIN 66131
- 2) following DIN ISO 787/XI, JIS K 5101/18 (not sieved)
- 3) following DIN ISO 787/II, ASTM D 280, JIS K 5101/21
- 4) following DIN 55 921, ASTM D 1208, JIS K 5101/23
- 5) following DIN ISO 787/IX, ASTM D 1208, JIS K 5101/24
- 6) following DIN ISO 787/XVIII, JIS K 5101/20
- based on material dried for 2 hours at 105°C
   based on material ignited for 2 hours at 1000°C
- 9) HCl content is a constituent of the ignition loss

Such pyrogenically prepared silicas are known. They are described, inter alia, in:

Winnacker-Küchler, Chemische Technologie, Volume 3 (1983), 4th edition, page 77 and

5 Ullmanns Encyklopädie der technischen Chemie, 4th edition (1982), Volume 21, page 462.

The pyrogenically prepared silicas are treated with a compound from the group  $(RO)_3SiC_nH_{2n+1}$ , wherein n= from 2 to 18 and R= alkyl, such as, for example, methyl, ethyl or the like.

The following compounds in particular can be used:

silane I  $(CH_3O)_3SiC_{16}H_{33}$  (hexadecyltrimethoxysilane)

silane II (CH<sub>3</sub>O)<sub>3</sub>SiC<sub>8</sub>H<sub>17</sub> (octyltrimethoxysilane).

The silicas that can be used according to the invention can
be prepared by placing the pyrogenically prepared silicas
in a mixer, spraying them, with intensive mixing,
optionally first with water and then with the compound
(organosilane) from the group (RO)<sub>3</sub>SiC<sub>n</sub>H<sub>2n+1</sub>, mixing for
from 15 to 30 minutes and then tempering at a temperature
of from 100 to 160°C for a period of from 1 to 3 hours,
followed by structure modification and/or optional milling.
A further tempering operation may optionally be carried out
after the structure modification and/or milling.

Structure modification can be carried out, for example,
using a ball mill or a continuously operating ball mill.
Milling can be carried out, for example, by means of an
air-jet mill or a pinned disk mill. Tempering can be
carried out batchwise, for example in a drying cabinet, or
continuously, for example in a fluid or fluidised bed.

Tempering can take place under protecting gas, for example nitrogen.

The water that is used can be acidified with an acid, for example hydrochloric acid, to a pH value of from 7 to 1.

The organosilane that is used can be dissolved in a solvent, such as, for example, ethanol.

5 Tempering can be carried out in a protecting gas atmosphere, such as, for example, under nitrogen.

The pyrogenically prepared silicas according to the invention silanised with silane I have the physicochemical data listed in Table 2 prior to structure modification:

Starting material	A 90	A 130	A 150	A 200	A 300	A 380	0X 20	тт 600
Average primary particle size [nm]	20	16	14	12.	7	7	40	40
BET surface area [m²/g]	40 - 90	60 - 130	75 - 150	100 - 200	150 - 300	200 - 380	20 – 50	100 - 250
Tamped density [g/l]	40 - 140	40 - 140	40 - 140	40 - 140	40 - 140	40 - 140	40 - 140	40 - 140
Loss on drying [%]	<b>2</b> 2	× 2		< 2	< 2	< 2	< 2	< 2
Ignition loss [%]	0.1 - 10	0.1 - 10	0.1 - 10	0.5 - 15	0.5 - 20	0.5 - 25	0.1 - 10	0.1 - 20
C content [%]	0.1 - 10	0.1 - 10	0.1 - 10	0.5 - 15	0.5 - 20	0.1 - 25	0.1 - 10	0.5 - 20
pH value	3.5 - 5.5	3.5 - 5.5	3.5 ~ 5.5	3.5 - 5.5	3.5 - 5.5	3.5 - 5.5	3.5 - 5.5	3.5 - 5.5

Table

## Examples

The pyrogenically prepared silicas that are used have the physicochemical data listed in Table 1.

The following compound of the general formula  $(RO)_3SiC_nH_{2n+1}$  is used as the organosilane:

(silane I)  $(CH_3O)_3SiC_{16}H_{33}$ 

The silica is placed in a mixer and sprayed, with intensive mixing, first with water and then with organosilane.

10 When the spraying is complete, mixing is carried out for a further 15 to 30 minutes, followed by tempering for from 1 to 3 hours at from 100 to 160°C. Tempering may also be carried out under protecting gas, for example nitrogen.

The individual reaction conditions can be taken from 15 Table 3.

The physicochemical data of the resulting silanised silicas are listed in Table 4.

Table 3

Tempering temperature (°C)	120	140	140	140	140	. 140
Tempering time (h)	2	2	2	2	2	2
Amount of ethanol (g/100 g	6	0	0	0	0	.0
Amount of water (g/100 g	0	0		2.5	1.25	1.25
Amount of silane (g/100 g	1	2.5	20	10	V	2.5
Silane	Silane I					
Aerosil	A 300	A 200				
Example	г	2	ю	ħ	5	9

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υ	
덥	I
Tab	İ
	•

Surface area Loss on drying Ignition loss (m²/g) (%)	1.8	2.5	12.7	7.1	3.4	2.5
Loss on drying (%)	0.4	0.3	9.0	9.0	9.0	0.7
Surface area (m²/g)	253	176	116	144	167	171
C content (%)	1.3	1.7	10.1	5.7	2.6	1.9
Tamped density (g/1)	. 50	49	89	72	52	51
pH value	4.3	4.4	9.4	4.5	4.7	4.5
Example	T	. 2	33.	4	5	9

# Preparation of the silicas used according to the invention:

The silidas, which can be prepared as described in EP 0 672 731, are subsequently subjected to structure modification by mechanical action and then optionally milled in a mill. Tempering may optionally be carried out after the structure modification and/or milling.

Structure modification can be carried out, for example, using a ball mill or a continuously operating ball mill.

Milling can be carried out, for example, by means of an air-jet mill or a pinned disk mill. Tempering can be carried out batchwise, for example in a drying cabinet, or continuously, for example in a fluid or fluidised bed. The tempering can take place under protecting gas, for example nitrogen.

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Table 5 Overview of the preparation of the comparison silicas and of the silicas according to the invention (Examples)

Description	Group attached to the surface	Structure modification	Milling after structure modification	Tempering after milling
Comparison silica 1	Hexadecylsilyl	No		
Comparison silica 2	Octylsilyl	No .	1	
Silicas 1	Hexadecylsilyl	Yes	No	No
Silicas 2	Octylsilyl	Yes	Yes	No
Silicas 3	Hexadecylsilyl	Yes	Yes	Yes
Silicas 4	Octylsilyl	Yes	No	Yes
Silicas 5	Octylsilyl	Yes	Yes	No
Silicas 6	Hexadecylsilyl	Yes	Yes	No
Silicas 7	Hexadecylsilyl	Yes	Yes	No
Silicas 8	Hexadecylsilyl	Yes	No	No
Silicas 9	Octylsilyl	Yes	Yes	No
Silicas 10	Octylsilyl	Yes	No	No
Silicas 11	Octylsilyl	Yes	Yes	No
Silicas 12	Octylsilyl	Yes	No	No

Table 6: Physicochemical data of the silicas according to the invention (Examples) and of the comparison silicas

Description	Tamped	Loss on	Idnition	pH value	Iquition   pH value   C content[%]	DBP	Specific surface
いっちょうけっちゅう	density	drving [8]	10ss [8]			adsorption	area according to
	[ fg/1]					[8]	BET [m2/g]
Comparison silica 1	57	0.5	1.8	4.6	1.2	302	195
Comparison silica 2	51	0.6	6.8	5.3	5.4	263	175
Silicas 1	137	0.7		4.9	1.3	217	193
Silicas 2	112	0.7	7.0	5.8	5.5	145	175
Silicas 3		0.7	2.3	5.1	1.3	228	176
Silicas 4		0.9	6.7	5.3	5.4	134	176
Silicas 5	114	0.5	7.1	0.9	5.4	142	175
Silicas 6	.	1.3	2.2	5.1	1.4	221	193
Gilicas 7		0.7	2.6		1.4	208	197
Silicas 8		1.1	2.3	5.8	1.4	182	195
Silicas 9	240	0.8	6.7	4.8	5.3	87	169
Silicas 10	322	0.3	6.9	0.9	5.3	not deter-	172
						minable	
Silicas 11	204	0.7	6.4	5.7	5.4	101	173
Silicas 12	276	0.3	9.9	9.9	5.3	not deter-	168
						minable	

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- (2) Silanised structure-modified silicas, which are characterised by groups attached to the surface, the groups being dimethylsilyl and/or monomethylsilyl, preferably dimethylsilyl, having the following physicochemical data:
- 5 BET surface area m²/g: 25 400
  Average primary particle size nm: 5 50
  pH value: 3-10
  Carbon content %: 0.1-10
  DBP number %: < 200
- 10 The silica that can be used according to the invention can have a tamped density of from 100 to 280 g/l, preferably from 100 to 240 g/l.

Pyrogenic silicas are known from Winnacker-Küchler Chemische Technologie, Volume 3 (1983), 4th edition,

15 page 77 and
Ullmanns Enzyklopädie der technischen Chemie, 4th edition
(1982), Volume 21, page 462.

Pyrogenic silicas are prepared especially by flame hydrolysis of vaporisable silicon compounds, such as, for example, SiCl<sub>4</sub>, or organic silicon compounds, such as trichloromethylsilane.

The invention also provides a process for the preparation of the silanised, structure-modified, pyrogenically prepared silicas according to the invention, which process is characterised in that pyrogenically prepared silica is treated in a known manner with dimethylchlorosilane and/or monomethyltrichlorosilane, the groups dimethylsilyl and/or monomethylsilyl being attached to the surface of the pyrogenic silica, subsequently subjected to structure modification and then optionally milled.

In an embodiment of the invention, tempering may be carried out following the structure modification and/or milling.

The silicas that can be used according to the invention can be prepared, for example, as follows:

The silicas, which can be prepared as described in DE 1 163 784, are subsequently subjected to structure modification by mechanical action and then optionally milled in a mill. Tempering may optionally be carried out following the structure modification and/or milling.

Structure modification can be carried out, for example, using a ball mill or a continuously operating ball mill.

10 Milling can be carried out, for example, by means of an air-jet mill or a pinned disk mill. Tempering can be carried out batchwise, for example in a drying cabinet, or continuously, for example in a fluid or fluidised bed. Tempering can be carried out under protecting gas, for example nitrogen.

## Examples

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# Preparation and physicochemical properties of the silicas

Preparation of the silicas that can be used according to the invention:

The silicas, which are prepared as described in DE 1 163 784, are subsequently subjected to structure modification by mechanical action and then optionally milled in a mill. Tempering may optionally be carried out following the structure modification and/or milling.

The individual parameters for the preparation of the individual silicas are listed in Table 7, and the physicochemical parameters are listed in Table 8.

Structure modification can be carried out, for example,
using a ball mill or a continuously operating ball mill.
Milling can be carried out, for example, by means of an
air-jet mill or a pinned disk mill. Tempering can be
carried out batchwise, for example in a drying cabinet, or
continuously, for example in a fluid or fluidised bed.
Tempering can be carried out under protecting gas, for
example nitrogen.

rable 7: Preparation of the silicas that can be used according to the invention

Description	Group attached to the surface	Structure modification	Milling after structure modification	Tempering after milling
Silie 1		Yes	МО	No
8i1is 2		Yes	No	NO
Silica 3			No .	NO
Stlica 4			No	ИО
Silica 5		Yes	Yes	No
Silica 6			No	No
Silica 7			Yes	No
Silica 8			Yes	хеs
Silica 9			Yes	Yes
Silica 10			Yes	No
Silica 11			Yes	No

Table 8: Physicochemical data of the silicas that can be used according to the invention

Description	Tempod density [g/1]	Loss on drying [%]	ignition loss [%]	enter Hg	pet value C content [%]	1889 adsomption [%]	Specific surface area according to BET [m²/g]
Silica 1	236	0.1	0.6	4.0	0.8	127	115
Silica 2	204	0.1	9.0	3.9	0.8	137	116
Silica 3	223	. 0.3	0.7	4.2	1.0	160	169
Silica 4	. 186	0.3	7.0	4.2	1.1	152	171
Silica 5	109	0.2	0.7	4.4	1.1	159	. 167
Silica 6	193	1.2	6.0	5.2	1.7	157	258
Silica 7	125	0.2	. 0.7	4.1	0.8	130	110
Silica 8	108	0.7	1.3	5.0	1.7		. 257
Silica 9	123	0.3	0.5	4.3	1.1	157	165
Silica 10	102	7.0	1.2	6.2	1.7	164	256
silica 11	160	0.2	2.0	- 4.0	0.8	132	115

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The silicas, which can be prepared as described in EP 0 672 731 or DE 1 163 784, are subsequently subjected to structure modification by mechanical action and then optionally milled in a mill. Tempering may optionally be carried out following the structure modification and/or milling. Structure modification can be carried out, for example, using a ball mill or a continuously operating ball mill. Milling can be carried out, for example, by means of an air-jet mill or a pinned disk mill. Tempering can be carried out batchwise, for example in a drying cabinet, or continuously, for example in a fluid or fluidised bed. Tempering can be carried out under protecting gas, for example nitrogen.

- (3) Low-structured pyrogenic metalloid and metallic oxides based on SiO<sub>2</sub>, Al<sub>2</sub>O<sub>3</sub>, SiO<sub>2</sub>/AlO<sub>3</sub>, TiO<sub>2</sub>, ZrO<sub>2</sub>, as described by EP 0 637 616.
- (4) Surface- and structure-modified products which are known from EP 0 808 880 A2, are already available commercially, such as, for example, AEROSIL® R 8200,
   AEROSIL® R 7200 (both Degussa), AEROSIL® R 972 W (Nippon Aerosil Corporation), WACKER HDK H 2000.

Surface- and structure-modified pyrogenic metalloid or metallic oxides are used according to the invention as flow improvers and anticaking agents. The way in which they act is described hereinbelow:

Materials having a low melting point, for example fats or waxes, acquire a soft surface as temperatures increase. The result is that the individual particles adhere to one another. By the addition of surface- and structure-modified pyrogenic silicas and metallic oxides, it is possible to coat the individual particles and separate them from one another. Caking of the particles is thus effectively

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prevented. In pharmacy (galenics), the prevention of sintering processes of pharmaceutical active ingredients, for example during pressing into tablets, also plays an important role.

5 Solid bridges between particles can form by recrystallisation and structure modification. In addition, during storage of mixtures of materials whose components are capable of reacting chemically with one another, reactions between those materials can occur, especially with the absorption of ambient moisture. Such reactions result in caking, which greatly reduces pourability. Providing such a mixture of materials with surface- and structure-modified pyrogenic silicas and metallic oxides prevents caking by drawing off the liquid film by suction, or by coating the particles and the spacer function associated therewith.

If a material consists of particles of different sizes, separation can readily occur because the larger particles move more rapidly than the smaller ones during transfer to a different container or during feeding. The addition of surface— and structure—modified pyrogenic silicas and metallic oxides effects coating of the particles, as a result of which a ball-bearing effect occurs. In addition, a certain standardisation of the particle size is achieved, so that separation is no longer to be feared. The shape of the particles present in a material is also of considerable importance for the pourability of materials. An uneven surface can result in increased friction and/or interlocking of the particles.

Many materials develop a film of moisture on the surface of their particles. During storage, the individual particles are in close contact. The surface tension of the liquid holds the particles together. If the material is soluble in water, recrystallisation and bridge formation may occur with variations in the atmospheric humidity. The addition

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of silica results in the film of moisture being drawn off by suction, the particles being surrounded completely by the silica. As a result, spatial separation occurs. If the material is hygroscopic, surface- and structure-modified pyrogenic silicas and metallic oxides are particularly effective.

In the case of pulverulent materials, VAN-DER-WAALS forces occur by the interaction of dipole moments of molecules.

In the case of non-conductive powders, an excess charge is
frequently observed, which leads in the case of antipolar
charging to forces of attraction. During the feeding of
powders through pipes or during mixing, sieving and milling
operations, that electrostatic charge always occurs.
Surface- and structure-modified pyrogenic silicas and
metallic oxides can reduce the electrostatic charge.

The surface- and structure-modified pyrogenic silicas and metallic oxides can be used according to the invention as flow regulators and anticaking agents in a concentration of from 0.001 to 50 wt.%, preferably from 0.01 to 10 wt.% and particularly preferably from 0.1 to 5 wt.%. The solids mixtures are prepared by mixing by means of known mixing devices. According to the invention, mixing devices that exert only low shear energy, such as, for example, gravity or ploughshare mixers, are suitable.

- Depending on the field of use, the solids mixtures can be processed further in different ways, for example introduction into capsules, dispersion, granulation, compacting, dissolution, milling, mixing, pelletisation, sifting, sieving, pressing into tablets, comminution, etc..
- The pulverulent materials prepared according to the invention with surface- and structure-modified pyrogenically prepared silicas and metallic oxides are distinguished by high flowability or fluidisability.

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The flowability is determined by numerous factors: the properties of the starting material (for example its grain size, chemical composition such as specific weight), the concentration and method of incorporation of the flow regulator, the atmospheric humidity, the storage time.

In order to determine the flowability of the pulverulent materials, siliconised glass discharge vessels having different outlet diameters may be used.

Table 9: Scheme for evaluating flowability using glass discharge vessels.

Vessel no. Rating number	Outlet width  Ø mm	Evaluation with still smooth passage of the powder
1	2.5	very good
2 .	5	good
3	8	on the whole good
4	12	adequate
5	18	poor
6		unsatisfactory (does not run through no. 5)

The height of those vessels is 90 mm, the inside diameter is 42 mm. Evaluation is made using the ratings 1 = very good flow behaviour, i.e. the test powder flows without stopping from glass discharge vessel no. 1 with the smallest outlet opening, to rating 6 = unsatisfactory flow behaviour, i.e. the powder does not even flow through glass discharge vessel no. 5 with the largest outlet opening. Table 9 shows discharge values and evaluation criteria. In

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general, the procedure is such that the powder test starts with glass discharge vessel no. 5 and the discharge vessels having the smaller openings are then tested in succession. The glass discharge vessel from which the powder just about flows without stopping is indicated.

A further method is measurement of the cone height of a powder. A metal sieve is secured at a distance of from 30 to 100 mm above a solid metal cylinder having a diameter of 50 mm and a height of about 80 mm. The distance between the metal sieve and the metal cylinder is governed by the flowability of the powder to be measured and should be slightly greater than the cone height of the powder of the test series to be tested that has the poorest pouring behaviour. The powder is shaken onto the sieve and slowly pressed through the sieve by hand with the aid of a plastics spatula. The powder that falls through forms a cone on the metal cylinder. Powder is passed through the sieve until a cone of geometrically uniform shape has formed on the cylinder. The sieve is then removed and the height of the cone is measured. From the height of the cone and the diameter of the metal cylinder it is possible to calculate the gradient of slope of the tested powder. Because the diameter of the cone is constant, the height of the cone can also be used as a direct measure of the pourability. Experience hitherto has shown that powders having cone heights of from 15 to 20 mm can be termed very readily pourable. Products having cone heights of from 50 to 70 mm have poor pouring behaviour. The values listed in Table 10 below are appropriate if value analysis is to be carried out according to different criteria or measuring methods.

Table 10

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Cone height mm	Assessment	•
	Description	Rating
< 20	very good	1
21 - 30	good	2
31 - 40	on the whole good	3
41 - 50	adequate	4
51 - 60	poor	5
> 60	unsatisfactory	6

The pulverulent materials according to the invention can be used in numerous applications and sectors, such as, for example, cosmetics, agriculture, foodstuffs, food supplements, pharmacy, animal feeds, etc..

There are listed hereinbelow examples of pulverulent materials which can be provided according to the invention with the surface- and structure-modified metalloid or metallic oxides as flow regulators:

covering powders, aminosulfonic acid, inorganic salts, aspirin, bath salts, brewer's yeast powder, lead oxides, lead and titanium dioxide, Carbowax 6000, cattle dust, cellulose powder, chilli powder, choline chloride powder, dragée production, fertilisers, egg powder (from egg yolk), egg powder (whole egg), iron sulfate heptahydrate, fat concentrates, fire-extinguishing powders, fish food, feeds, spice mixtures, foundry auxiliaries, gypsum, guar gum, urea (cryst.), urea (prilled), domestic fertilisers,

hexamethylenetetramine, HVP (hydrolized vegetable powders), industrial salts, instant drink powders, coffee powder, coffee whiteners, cocoa powder, potato starch, cheese (grated cheese), cheese powder (parmesan), adhesive powders, garlic powder, cooking salt, cosm. covering 5 powders, plastics films, skimmed milk powder, corn starch, malt powder, molasses, melamine resin powder, methionine, milk substitute, milk powder, mineral mixtures, whey powder, monoammonium phosphate, sodium hydrogen carbonate, 10 sodium hydrogen sulfate, sodium perborate, sodium propionate, sheets of paper, paprika powder, pesticides, plant-protecting granules, plant-protecting dusts, polyethylene powder, powdered sugar, pigments, pickling salt, polymers, proxyphylline, powder, powdered rubber, 15 powder coatings, PVC powders, rice starch, roast sugar, inorganic salts in general, scouring powders, sulfur, soap powders, silver halides, sintered metal powders, table salt, disintegrators, wettable powders (plant protection), fruit powders, trace element pre-mixes, S-PVC powders, soup 20 powders, tomato powders, toners, toilet cleaners, powder type fire extinguishers, vitamin pre-mixes, detergents, anhydrous citric acid, fluidised sintered powders, zirconium oxide, citrus powder, onion powder, sugars.

An example of a pulverulent material according to the invention and the use according to the invention of the surface-modified and structure-modified pyrogenically prepared metalloid and metallic oxides is the improvement to the fluidisability of fire-extinguishing powders. Fire-extinguishing powders should be readily fluidisable even after a prolonged storage period and must not cake so that, in the event of a fire, problem-free spraying from the extinguisher is ensured. To that end, small amounts of a flow improver are added thereto according to the prior art.

35 Such a flow improver may be, for example, conventional

pyrogenic or precipitated, hydrophobic or hydrophilic silicas, silicates or special oxides. The best results according to the prior art are achieved with hydrophobic pyrogenic silicas such as, for example, Aerosil® R972. Nevertheless, the fluidisability achieved therewith is

5 Nevertheless, the fluidisability achieved therewith is still not sufficient.

Surprisingly, the use of the structure-modified aerosils according to the invention led to a marked improvement in the fluidisability and a marked reduction in the tendency to caking compared with the prior art.

## Table 11:

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Overview of the preparation of the comparison silicas and of the silicas that can be used according to the invention

Description	Group attached to the surface	Structure modifica- tion	Milling after structure modification	Tempering after milling
Example 1 R 974	Dimethylsilyl	Yes	No	No
Example 2 R 974	Dimethylsilyl	Yes	Yes	No
Example 3 R 974	Dimethylsilyl	Yes	Yes	Yes
Example 4 R 972	Dimethylsilyl	Yes	. No	No
Example 5 R 972	Dimethylsilyl	Yes	Yes	No

The physicochemical data of the silicas are listed in Table 12.

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Description	Tamped density [g/1]	Loss on drying [%]	Ignition loss [%]	pH value	C content [%]	DBP adsorption [%]	Specific surface area according to BET [m²/g]
Aerosil R 972 (comparison silica)	64	0.1	0.5	4.0	0.8	243	113
Aerosil R 973 (comparison silica)	. 67	0.5	9.0	4.8	1.0	256	165
Aerosil 200 (comparison silica)	56	1.1	0.4	4.0	0	nđ	206
Aerosil R 972 W (according to the invention)				4.38	.0.78		108
Example 1 (according to the invention)	186	0.3	0.7	4.2	1.1	152	171
Example 2 (according to the invention)	109	0.2	0.7	4.4	1.1	159	167
Example 3 (according to the invention)	123	0.3	0.5	4.3	H H	157	165
Example 4 (according to the invention)	204	0.1	9.0	3.9	0.8	137	116
Example 5 (according to the invention)	125	0.2	0.7	4.1	0.8	130	110
Aerosil R 7200 (according to the invention)	258	8.0	8.1	4.4	5.0	nđ	155
Aerosil R 8200 (according to the invention)	175	0.3	3.2	5.7	3.1	nđ	161
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Fire-extinguishing powder (preparation of the samples)

495 g of the fire-extinguishing powder and 5 g of the silica are weighed into a 1000 ml screw-top jar and the cover is closed. The materials are mixed for 30 minutes at moderate speed using a Turbula mixer.

The fluidisability of a fire-extinguishing powder is measured by means of a fluidising apparatus consisting of two 750 ml Erlenmeyer flasks which are connected together. To that end, 500 g of extinguishing powder mixture (fireextinguishing agent based on ammonium salts and mineral additives) are weighed into one of the Erlenmeyer flasks, and the second Erlenmeyer flask is fastened to the first by means of a sealing ring and a connecting sleeve. The apparatus, which resembles an hour-glass, is clamped in a 15 rotatable holder on a stand, and the powder is allowed to settle for 30 minutes. The apparatus is then rotated through 180° so that the extinguishing powder flows from one flask into the other. As soon as the extinguishing powder has flowed out of one flask, it is again rotated 20 through 180°. 30 passes are carried out in immediate succession, and the time at 10, 20 and 30 passes is determined.

The time for 10, 20 and 30 passes is recorded and the differences in the times are calculated according to the calculation below. The times for the first 10, the second 10 and the third 10 passes are thus determined. Shorter times mean better fluidisability.

 $t_1$  = time for 10 passes

 $t_2$  = time for 20 passes

30  $t_3 = time for 30 passes$ 

 $\Delta t_1 = t_1 = time for the first 10 passes$ 

 $\Delta t_2 = t_2 - t_1 = time for the second 10 passes$ 

 $\Delta t_3 = t_3 - t_2 = time for the third 10 passes$ 

Table 13

Example	Type of modification	t <sub>1</sub> [s]	t <sub>2</sub> [s]	t <sub>3</sub> [s]	Δt <sub>1</sub> [s]	Δt <sub>2</sub>	Δt <sub>3</sub> [s]
Aerosil R974	None	350	703	1040	350	353	337
1	Structure modification	343	407	476	343	64	69
2	Structure modification and milling	319	405	480	319	86	75
3	Structure modification, milling and tempering	197	265	333	197	68	68

Table 13 shows the fluidisability of fire-extinguishing powders. Shorter times mean an improvement in the fluidisability.

A markedly improved fluidisability of the fireextinguishing powder compared with the prior art is found with structure-modified aerosils, especially after more than 10 passes.

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## Determination of the tendency to caking

#### Test method 1

The tendency to caking is tested by placing 50.0 g of test material into a 100 ml glass beaker and placing the beaker in a 600 ml glass beaker containing a layer of water of about 2 cm. The glass beaker is closed tightly with aluminium foil and stored in a drying cabinet for 15 hours at 80°C. The 100 ml glass beaker is then removed from the 600 ml glass beaker and the test material is dried for 15 hours at 80°C. The sample is then sieved manually through a 0.5 mm sieve and the residue is determined; the smaller the residue, the better the prevention of caking. The appearance of the sample is also assessed.

Table 14: Determination of the tendency of fireextinguishing powders to caking

Example	Type of modification	Sieve residue [g]	Visual assessment
Aerosil R974	None	51.0	Slighly moist, very pronounced caking, many agglomerates, phosphate fusion occurs
2	Structure modification and milling	6.8	Still as dry, disintegrates to the original degree of dispersion when rolled and shaken, many agglomerates, no phosphate fusion occurs
3	Structure modification, milling and tempering	9.2	Still as dry, disintegrates to the original degree of dispersion when rolled and shaken, many agglomerates, no phosphate fusion occurs

### 5 Test method 2

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20.0 g of sample material are introduced into a Petri dish having a diameter of 5 cm, and the dish is stored in a desiccator over saturated ammonium sulfate solution for 24 hours at room temperature and 80 % atmospheric humidity. The appearance of the powder is then assessed.

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Table 15: Determination of the tendency of fireextinguishing powders to caking

Example	Type of modification	Sieve residue [g]	Visual assessment
Aerosil R 974	None	51.0	Slightly moist, very pronounced caking, many agglomerates, phosphate fusion occurs
2	Structure modification and milling	6.8	Still as dry, disintegrates to the original degree of dispersion when rolled and shaken, many agglomerates, no phosphate fusion occurs
3	Structure modification, milling and tempering	9.2	Still as dry, disintegrates to the original degree of dispersion when rolled and shaken, many agglomerates, no phosphate fusion occurs

<u>Table 16:</u> Tendency of fire-extinguishing powders to caking (test method 2)

Example	Type of modification	Visual assessment
Aerosil R 974	None	Moist, few agglomerates, the original degree of dispersion is not achieved when shaken, no phosphate fusion occurs
2	Structure modification and milling	Slightly moist, no agglomerates, disintegrates to the original degree of dispersion when rolled and shaken, no phosphate fusion occurs
3	Structure modification, milling and tempering	Slightly moist, no agglomerates, disintegrates to the original degree of dispersion when rolled and shaken, no phosphate fusion occurs

According to both test methods, there is a markedly reduced tendency to caking compared with the prior art when fire-extinguishing powders are provided with the structure-modified aerosils used according to the invention.

## Example 2 (pharmaceutical excipient mixtures)

Determination of the agglomerate content and the flowability of mixtures with microcrystalline cellulose for pharmaceutical applications

198.0 g of Avicel PH101 and in each case 2.0 g of silica ' and the products from the Examples were pre-mixed manually in 1 1 wide-necked flasks. The powder mixtures were then transferred to a set of sieves (stainless steel) consisting of two sieves having mesh widths of 0.71 mm (upper) and 0.315 mm (lower) and a closed sieve bottom. The mixtures were sieved, by manual shaking, until the amount of solid on the two sieves did not decrease further. The two sieves and the sieve bottom, whose empty weights had first been determined, were then weighed in order to determine the 15 amount of silica agglomerates which were too coarse to pass through the 0.71 mm or 0.315 mm sieve. The three sieved fractions of a mixture were then again combined in the 1 l three-necked flask and mixed in a gravity mixer (Turbula) for 5 minutes at 42 revolutions per minute. The flow rating 20 and the cone height of the mixture were then determined, before the content of the sieved fractions was again determined as described above. The fractions were again combined and mixed for a further 55 minutes. The flow rating, the cone height and the amount of the sieved 25 fractions were again determined.

The results of the tests are summarised in Table 17 below.

Table 17:

	Comparison ex	amples		Examples		
Mixing time		Aerosil 200	Aerosil R972	Aerosil R 972W (NAC)	Example 4	Example 5
	Sieve residue 0 min 0.71 mm	64.0 %	46.0%	0	0	5.5 %
0 min	Sieve residue 0 min 0.315 mm	32.5 %	20.5 %	1.0 %	8.0 %	9.5 %
	Sieve residue 0 min Total	96.5' %	66.5 %	1.0 %	8.0 %	15.0 %
·	Sieve residue 5 min 0.71 mm	58.5 %	39.5 %	0	4.5 %	10.0 %
	Sieve residue 5 min 0.315	28.0 %	18.0 %	1.5 %	10.5	16.5 %
5 min	Sieve residue 5 min Total	86.5	57.5 %	1.5 %	15.0 %	26.5 %
	Flow rating	6	4	3	2	3
	Cone height (cm)	2.5	2.0	1.75	1.9	1.9
	Sieve residue 60 min 0.71 mm	33.5 %	32.0 %	0	4.5 %	2.5 %
60 min	Sieve residue 60 min 0.315 mm	15.0 %	18.0 %	1.5 %	12.0 %	8.5 %
	Sieve residue 60 min Total	48.5 %	50.0 %	1.5 %	15.5 %	11.0 %
	Flow rating	4.5	3.5	2	2	2.5
	Cone height	2.3	1.9	1.95	1.9	1.9

		Examples			····	··
Mixing time		Example	Example 2	Example 7	Aerosil R7200	Aerosil R8200
	Sieve residue 0.71 mm	10.5 %	1.0 %	4.5 %	0.05 %	14.0 %
0 min	Sieve residue 0.315 mm	12.5 %	7.0 %	4.5 %	3.0 %	15.0 %
	Sieve residue Total	23.0 %	8.0 %	9.0 %	3.5 %	29.0 %
	Sieve residue 0.71 mm	10.5 %	6.0 %	5.5 %	2.0 %	12.5 %
5 min	Sieve residue 0.315 mm	15.0 %	5,0 %	5.5 %	2.0 %	16.5 %
	Sieve residue Total	25.5 %	11.0%	11.0 %	4.0 %	29.0 %
	Flow rating	3	3	3	4	2 ·
	Cone height	1.95	1.85	1.9	1.9	1.8
	Sieve residue 0.71 mm	5.0 %	1.0 %	0.0 %	1.0 %	1.0 %
60 min	Sieve residue 0.315 mm	8.0 %	2.0 %	5.0 %	1.5 %	1.5 %
	Sieve residue Total	13.0 %	3.0 %	5.0 %	2.5 %	2.5 %
	Flow rating	3	3	4	3	1
	Cone height	1.85	2	1.9	1.9	1.8

The tests clearly show that the powder mixtures according to the invention containing the surface- and structure-modified pyrogenic silicas Aerosil R 972 W, the products of the Examples and Aerosil 200 and Aerosil R 972 have a markedly smaller sieve residue and hence agglomerate content than the comparison products Aerosil 200 and Aerosil R 972. In addition, the maximum flowability is achieved after a mixing time of only 5 minutes whereas, when AEROSIL 200 and AEROSIL R 972 are used, the maximum flowability achievable with those materials is reached only after 60 minutes.

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### Patent claims:

- 1. Pulverulent materials and mixtures thereof, characterised in that they contain one or more surface-modified and structure-modified pyrogenically prepared metalloid or metallic oxides.
- 2. Method of improving the flowability of pulverulent materials and mixtures thereof, characterised in that there are added to the pulverulent materials and mixtures thereof one or more surface-modified and structure-modified pyrogenically prepared metalloid or metallic oxides.
- 3. Use of surface-modified and structure-modified pyrogenically prepared metalloid or metallic oxide for improving the flowability of pulverulent materials and mixtures thereof.

# INTERNATIONAL SEARCH REPORT

Internation	pplication No
PCT	03/11053

			03/11033
a. classi IPC 7	FICATION OF SUBJECT MATTER B01J2/30		
According to	International Patent Classification (IPC) or to both national classific	ation and IPC	
B. FIELDS	SEARCHED		
Minimum do IPC 7	cumentation searched (classification system followed by classification $B01J-A62D$	on symbols)	
	ion searched other than minimum documentation to the extent that s		
EPO-In	ata base consulted during the International search (name of data ba	se and, where pradical,	search terms used)
C. DOCUME	ENTS CONSIDERED TO BE RELEVANT		
Category °	Citation of document, with indication, where appropriate, of the rel	evant passages	Relevant to claim No.
X	US 5 959 005 A (HENNIG THOMAS ET 28 September 1999 (1999-09-28) column 1, line 53 - line 56; clait EP 0 808 880 A 26 November 1997 (1997-11-26) cited in the application	•	1-3
	ner documents are listed in the continuation of box C.	χ Patent family n	nembers are listed in annex.
'A' docume consid 'E' earlier of filing d 'L' docume which citation 'O' docume other of the results of the resu	nt which may throw doubts on priority claim(s) or is cited to establish the publication date of another in or other special reason (as specified) and referring to an oral disclosure, use, exhibition or neans ent published prior to the international filing date but	or priority date and cited to understand invention  "X" document of particul cannot be consider involve an inventive 'Y' document of particul cannot be consider document is combi	ished after the international filing date not in conflict with the application but it the principle or theory underlying the lar relevance; the claimed invention ed novel or cannot be considered to estep when the document is taken alone lar relevance; the claimed invention ed to involve an inventive step when the ned with one or more other such documention being obvious to a person skilled of the same patent family
Date of the	actual completion of the international search	Date of mailing of ti	ne International search report
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tion on patent family members

Internation	pplication No	
PCT.	03/11053	

Patent document cited in search report		Publication date		Patent family member(s)	Publication date
US 5959005	A	28-09-1999	DE CA CN DE EP JP KR	19616781 A1 2203726 A1 1167729 A ,B 59709069 D1 0808880 A2 10087317 A 260325 B1	06-11-1997 26-10-1997 17-12-1997 13-02-2003 26-11-1997 07-04-1998 01-07-2000